

# Differential impact of lactose/lactase phenotype on colonic microflora

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**BACKGROUND:** The ability to digest lactose divides the world's population into two phenotypes that may be risk variability markers for several diseases. Prebiotic effects likely favour lactose maldigesters who experience lactose spilling into their colon.

**OBJECTIVE:** To evaluate the effects of fixed-dose lactose solutions on fecal bifidobacteria and lactobacilli in digesters and maldigesters, and to determine whether the concept of a difference in ability to digest lactose is supported.

**METHODS:** A four-week study was performed in 23 lactose maldigesters and 18 digesters. Following two weeks of dairy food withdrawal, subjects ingested 25 g of lactose twice a day for two weeks. Stool bifidobacteria and lactobacilli counts pre- and postintervention were measured as the primary outcome. For secondary outcomes, total anaerobes, Enterobacteriaceae, beta-galactosidase and N-acetyl-beta-D-glucosaminidase activity in stool, as well as breath hydrogen and symptoms following lactose challenge tests, were measured.

**RESULTS:** Lactose maldigesters had a mean change difference (0.72 log<sub>10</sub> colony forming units/g stool; P=0.04) in bifidobacteria counts compared with lactose digesters. Lactobacilli counts were increased, but not significantly. Nevertheless, reduced breath hydrogen after lactose ingestion correlated with lactobacilli (r=-0.5; P<0.001). Reduced total breath hydrogen and symptom scores together, with a rise in fecal enzymes after intervention, were appropriate, but not significant.

**CONCLUSIONS:** Despite failure to achieve full colonic adaptation, the present study provided evidence for a differential impact of lactose on microflora depending on genetic lactase status. A prebiotic effect was evident in lactose maldigesters but not in lactose digesters. This may play a role in modifying the mechanisms of certain disease risks related to dairy food consumption between the two phenotypes.

**Key Words:** Fecal microflora; Lactose; Prebiotic

The ability (lactase persistent [LP]) or inability (lactase nonpersistent [LNP]) to digest lactose in adulthood is the most common known genetic polymorphism that divides humans into a dichotomous phenotype. The near-total loss of intestinal lactase by mid-childhood in the majority of LNP populations forces an altered pattern of food consumption and handling of lactose by this group (1,2). Because of symptoms incurred by intermittent consumption of lactose (mostly dairy foods [DFs]), lower quantities are generally consumed by LNP

## L'effet différentiel du phénotype de lactose-lactase sur la microflore colique

**HISTORIQUE :** La capacité de digérer le lactose divise la population mondiale en deux phénotypes qui peuvent risquer les marqueurs de variabilité de plusieurs maladies. Les effets prébiotiques favorisent probablement les personnes qui digèrent mal le lactose et subissent un épanchement de lactose dans le colon.

**OBJECTIF :** Évaluer les effets de solutions de lactose à doses fixes sur les bifidobactéries et les lactobacilles fécales chez les personnes qui digèrent bien ou mal le lactose et déterminer si le concept d'une différence de capacité de digérer le lactose est étayé.

**MÉTHODOLOGIE :** Les chercheurs ont effectué une étude de quatre semaines chez 23 personnes qui digèrent mal le lactose et 18 personnes qui le digèrent bien. Après deux semaines de sevrage des produits laitiers, les sujets ont ingéré 25 g de lactose deux fois par jour pendant deux semaines. Comme issue primaire, les chercheurs ont mesuré la numération de bifidobactéries et de lactobacilles dans les selles avant et après l'intervention. Comme issue secondaire, ils ont mesuré l'activité totale des anaérobies, des entérobactériacées, des bêta-galactosidases et des N-acétyl-bêta-D-glucosaminidases dans les selles, de même que l'hydrogène dans l'haleine et les symptômes après des tests de provocation par le lactose.

**RÉSULTATS :** Les personnes qui digèrent mal le lactose ont une différence de changement moyenne (0,72 log<sub>10</sub> unités formatrices de colonies/g de selle; P=0,04) dans les bifidobactéries par rapport à celles qui le digèrent bien. Les numérations de lactobacilles augmentaient, mais pas de manière significative. Néanmoins, une diminution d'hydrogène dans l'haleine après l'ingestion de lactose était corrélée avec les lactobacilles (r=-0,5; P<0,001). L'association d'une diminution d'hydrogène total dans l'haleine et des indices des symptômes, accompagnée d'une augmentation des enzymes fécaux après l'intervention, était appropriée, mais non significative.

**CONCLUSIONS :** Malgré l'échec à obtenir une adaptation colique complète, la présente étude a fourni des données probantes démontrant un effet différentiel du lactose sur la microflore d'après le statut génétique de la lactase. Un effet prébiotique était évident chez les personnes qui digèrent mal le lactose, mais pas chez celles qui le digèrent bien. Ce phénomène peut contribuer à modifier les mécanismes de certains risques de maladie liés à la consommation de produits laitiers entre les deux phénotypes.

populations (3,4). However, regular consumption of lactose may lead to bacterial metabolism of nondigested lactose that spills into the lower intestine (prebiotic effect) leading to colonic adaptation, decreased symptoms and further increased intake and tolerability (5-8).

The overwhelming concentration of research on improving symptoms of lactose intolerance in LNP subjects may mask the significance of bacterial colonic adaptation. Over the past few decades, several authors have postulated that there are

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was inadvertently inappropriately recorded. On completion of this period, stool and lactose challenge tests were repeated.

#### Lactase genetic tests

Blood drawn during the first phase of the study was used to determine the predominant genetic polymorphism in the lactase promoter region that was analyzed. The polymorphism (C/T-13910) associates completely with the promoter region of the LP/LNP gene in the majority of European populations (17,18). DNA was prepared using a commercially available DNA isolation kit (Gentra Systems, USA). A real-time polymerase chain reaction assay based on fluorescence resonance energy transfer (19,20) was used. The LightCycler DNA Master Hybridization Probes kit (Roche Diagnostics, Germany) was used for analysis of the C/T genetic polymorphism. The C/C genotype of this polymorphism is associated with LNP individuals, the T/T genotype is associated with LP individuals and the heterozygote C/T is also considered to be LP, but with a reduced measured level of intestinal lactase (17). As such, the cohort was classified into two phenotypes: lactose digester (TT and CT) and maldigester (CC).

#### BH<sub>2</sub> test

The BH<sub>2</sub> test is the only clinical test that physiologically reflects metabolic and/or bacterial changes to dietary interventions and assesses colonic adaptation (5,21). It was included in the present study primarily as a measure of the adaptation process. Clinically, colonic adaptation includes improved symptoms, a measurable decrease in BH<sub>2</sub> and an increase in fecal beta (β)-galactosidase (see below) on rechallenge after continued regular intake of lactose (5). Generally, the changes in BH<sub>2</sub> following intervention are statistically significant.

Briefly, BH<sub>2</sub> was measured in parts per million (ppm) using a validated hand-held hydrogen chemical sensor (EC60 gastrolyzer, Bedfont Scientific Ltd, United Kingdom) (22,23). Following a baseline measurement, subjects ingested 50 g of lactose mixed in water. BH<sub>2</sub> was then remeasured at 15 min, 30 min, 45 min, 60 min, 75 min and 90 min intervals following ingestion, and every 30 min thereafter for a total of 270 min (4.5 h). The baseline value was subtracted from readings recorded at each subsequent time interval. In general, an acceptable baseline value is 20 ppm or lower. A definite positive value is defined as more than 20 ppm above baseline at any time point (21). Results of each BH<sub>2</sub> measurement were summed to obtain a value for total breath hydrogen (TBH<sub>2</sub>).

#### Total symptom score

Symptoms were recorded on a four-point Likert scale at baseline and at 30 min intervals following ingestion of 50 g of lactose for 240 min. Bloating, gas and cramps were assigned a score of 0 if there were no symptoms, 1 for mild symptoms, 2 for moderate symptoms and 3 for severe symptoms. Diarrhea was scored as 0 (none) or 1 (present). The total symptom score (TSS) was obtained by summing all scores at each time point; the minimum possible score was 0, with a maximum possible score of 90 ( $[9 \times 3 \times 3] + 9 \times 1$ ).

#### Stool bacterial counts

Participants encountered no problems with successfully providing stool samples during each visit. Samples were coded, placed on ice and were shipped to the microbiology laboratory

(McGill University, Macdonald site, Montreal, Quebec) within 2 h. The samples were analyzed on the same day. No samples were sent on weekends.

A slightly modified method based on a previous report (24) on fecal bacteria testing was used. For quantification of total bifidobacteria, total lactobacilli, total anaerobes (bifidobacteria, lactobacilli, bacteroides and clostridia) and total enterobacteria, the fresh fecal samples (corrected to 100 g) were introduced into an anaerobic jar (BBL GasPak, Becton Dickinson, USA) containing a gas mixture. A 1.0 g sample of feces was removed and homogenized in 4 mL of prepared brain heart infusion broth supplemented with yeast extract and 5-cysteine hydrochloride. A serial dilution was made and subsequently plated onto Bifidobacterium (Beerens), Lactobacillus (Rogosa) agar, and Enterobacteriaceae (MacConkey) plates. The counts (colony forming units [CFU]) of total culturable anaerobes were enumerated on reinforced Clostridial medium agar (Fisher Scientific, USA). The plates were incubated anaerobically for two to three days at 37°C, with the exception of the MacConkey plates, which were incubated aerobically overnight at 37°C. All bacterial counts are reported as CFU/g of stool. The primary changes of interest were focused on bifidobacteria and lactobacilli because they are known to proliferate in vitro with regular lactose challenge; however, all four bacterial strains were analyzed and compared.

#### Stool enzyme activity

The bacterial lactase enzymes β-D-galactosidase (β-gal) and N-acetyl-β-D-glucosaminidase (NAG) were assessed as measures of stool enzyme activity. β-gal is a ubiquitous bacterial enzyme analogous to mammalian intestinal lactase, which increases with bacterial colonic adaptation (16,25). Measurement of fecal β-gal was performed using the O-nitrophenyl-β-D-galactopyranoside (ONPG) method (26). Briefly, 20 μL of stool in buffer was added to 480 μg of ONPG in sodium phosphate buffer (pH 7.0). The reaction was allowed to proceed at 45°C for 10 min. Sodium carbonate (1 M) was added to stop the reaction. Optical density at 420 nm was subsequently read and β-gal activity was reported as units/g of stool.

NAG is also a ubiquitous bacterial enzyme involved in mucus digestion and may exert bactericidal effects (27-29). Although NAG has not been previously evaluated in models of colonic adaptation, it is hypothesized that it may be a marker of bacterial response to lactose because of its role in mucus digestion. NAG was analyzed using a buffered sample of 0.1 mL of stool added to 1 mL of substrate. After incubation at 37°C for 30 min with 0.5 mL of sodium carbonate buffer to stop the reaction, the optical density was read at 405 nm (26), with NAG activity reported as units/mg of stool.

#### Statistical analysis

The demographics of the population are described using means and SDs for continuous variables and percentages with standard errors for categorical variables. Normally distributed data were obtained by log<sub>10</sub> transforming the bacterial CFU/g of stool and enzyme activity.

The primary objective was to determine whether lactose consumption increases bifidobacteria and lactobacilli counts differentially in LNP subjects versus LP subjects. The respective change in scores (ie, follow-up minus baseline) were calculated and reported as mean change with 95% CIs and P values. The change in

**TABLE 1**  
**Demographics**

	Dropouts (n=16)	Participating (genotype*)	
		LNP (CC) (n=23)	LP (CT/TT) (n=18)
Male sex, n (%)	8 (50)	7 (30)	8 (44)
Race, n (%)			
African	3 (19)	5 (22)	2 (11)
Asian	1 (6)	8 (35)	0 (0)
Caucasian	12 (75)	10 (43)	16 (89)
Genetics, n (%)			
CC	7 (44)	23 (100)	0 (0)
CT	6 (38)	—	10 (56)
TT	3 (19)	—	8 (44)
Age <sup>†</sup>	29.1±8.1	32.1±8.7	26.3±6.5
Body mass index <sup>†</sup> , kg/m <sup>2</sup>	22.7±3.1	23.8±4.3	22.8±2.4
Daily lactose intake at study entry <sup>‡</sup> , g/day	16.5 (0–60.1)	11.0 (4.5–36.9)	9.5 (0.4–26.7)

Categorical variables are shown as a per cent of total within that group. \*Refers to the polymorphism at position 13910 of the lactase promoter region; <sup>†</sup>Data for continuous measures are presented as mean ± SD; <sup>‡</sup>Data presented as median (interquartile range). Daily lactose intake is based on responses to a three-day recall questionnaire. LNP Lactase nonpersistent; LP Lactase persistent

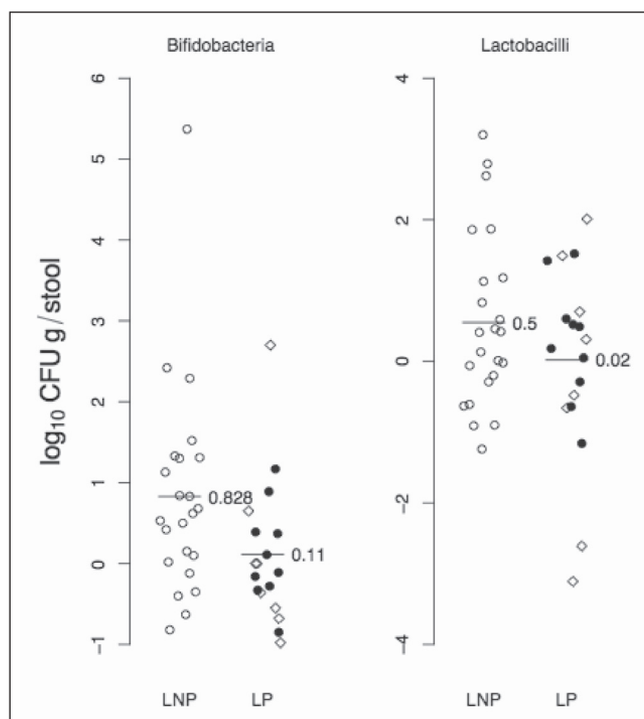
scores between follow-up and baseline for TBH<sub>2</sub>, TSS, log<sub>10</sub>(CFU/g stool) for Enterobacteriaceae and total anaerobic bacteria, log<sub>10</sub>(g/stool) β-gal and log<sub>10</sub>(units/mg stool) NAG are also described.

P<0.05 was considered to be statistically significant and P<0.1 was considered to be a trend. Because of the nature of the experiment, the analysis was limited to subjects who were measured at both baseline and follow-up. For missing BH<sub>2</sub> values and TSS at any particular time point, data were assigned using the following algorithm (which should bias the results toward the null hypothesis): if data were available for time points immediately before and following the time point with missing data, the average of the two was used. If there were no data before or following the missing data time point (eg, baseline missing or 270 min value missing), the average of the scores for that time point from the other visits (for this scenario, data from the visit four weeks before baseline were included) were used. For missing data regarding bacterial counts and enzyme analysis, the average score of the other visits was used. A sensitivity analysis for subjects with no missing data was also performed.

### Sample size

Based on a previous study using 15 LNP and 15 LP subjects (30), the two exposure groups (LNP and LP) were expected to be of near equal size. Sample sizes were calculated setting alpha at 0.05 and power at 0.8.

The calculated SD of change for lactobacilli over time in the study by Ito and Kimura (31) was 0.79 (log<sub>10</sub> units) using P=0.05 (the reported P value was less than 0.05; the calculation used in the present study represents a conservative approach when only pre and post SDs are provided, as in the Ito and Kimura study). A similar SD of change was assumed for the other primary outcome bifidobacteria count. There are no data to determine what a clinically relevant difference in bacterial counts should be. Therefore, a three-fold difference (0.48 log<sub>10</sub> units) was arbitrarily chosen to be clinically



**Figure 2** Scatter plot showing the change in the primary outcomes: bifidobacteria and lactobacilli counts (log<sub>10</sub> [colony forming units (CFU)/g stool]) between test 3 and test 2. Lactase nonpersistent (LNP) subjects are represented by open circles (genotype CC). Lactase persistent (LP) subjects are represented by solid circles (genotype TT) and open diamonds (genotype CT). In addition, the mean change for the respective LNP/LP groups is shown with a line and the exact value. No comparisons were made with test 1 (baseline)

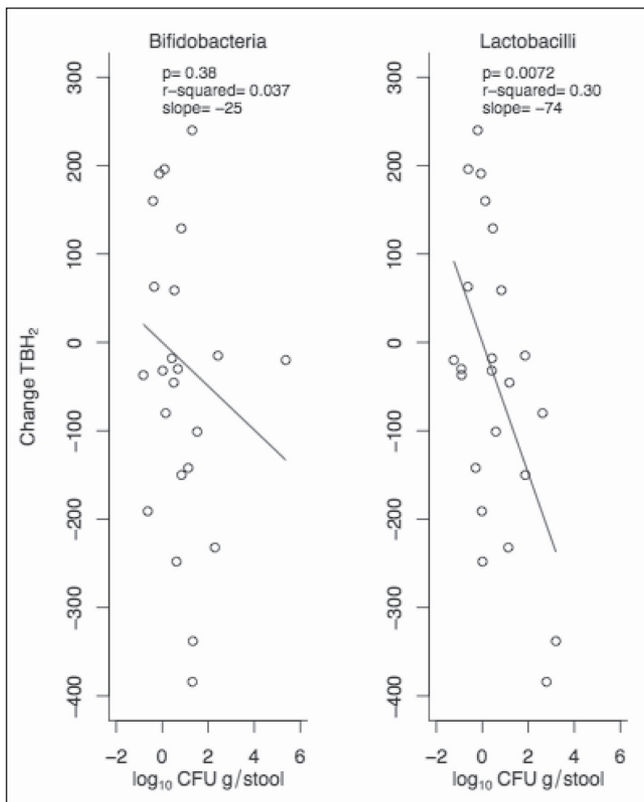
relevant. It was estimated that 45 participants per group would be required to detect a statistically significant difference.

## RESULTS

Participants were classified as LP or LNP based on the genetic lactase test. Although the study had initially intended to recruit 46 subjects per group, only a total of 57 were recruited because of logistical reasons. From the initial 57 participants, 16 did not complete the study (seven LNP subjects and nine LP subjects), leaving 41 with data for baseline and follow-up visits. The demographic characteristics of subjects who dropped out and participated are shown in Table 1. The distribution of participants and reasons for leaving the study are shown in Figure 1.

A scatter plot showing the mean change in primary outcomes (ie, bifidobacteria and lactobacilli counts) between tests 3 and 2 stratified according to LNP status is shown in Figure 2. The difference in the mean change in bifidobacteria counts (log<sub>10</sub> CFU/g stool) for LNP subjects (genotype CC) versus LP subjects (genotypes TT and CT) was 0.72 (95% CI 0.03 to 1.04; P=0.04).

As an exploratory posthoc analysis, the change in bifidobacteria counts were dichotomized to being positive based on thresholds of a 1log unit or greater change, and a 0.5log unit change. Of the LNP subjects, eight of 23 (35%) had a 1log unit change from baseline for bifidobacteria versus two of 18 (11%)



**Figure 3** Scatter plot showing the change in total breath hydrogen (TBH<sub>2</sub>) between test 3 and test 2, with the change in bacterial counts (colony forming units [CFU]) for the primary outcomes for bifidobacteria (**left panel**) and lactobacilli (**right panel**) in lactase nonpersistent subjects only (lactase persistent subjects have almost no TBH<sub>2</sub> to begin with). The P value, r<sup>2</sup> and slope are indicated in each panel. No comparisons were made with test 1 (baseline)

for LP subjects ( $\chi^2=1.92$ ,  $P=0.14$  [Fisher's exact test]). When a threshold of 0.5log units or greater was used, there were 14 of 23 (61%) LNP subjects with positive responses and four of 18 (22%) LP subjects ( $\chi^2=4.65$ ,  $P=0.03$  [Fisher's exact test]).

In a similar analysis for lactobacilli, the difference in the mean change was 0.53log<sub>10</sub> CFU/g stool (95%CI -0.30 to 1.36;  $P=0.20$ ). At a threshold of 1log unit or greater change from baseline, there were seven of 23 (30%) positive responses in LNP subjects and four of 18 (22%) positive responses in LP subjects ( $\chi^2=0.05$ ,  $P=0.73$  [Fisher's exact test]); the corresponding figures for a threshold of 0.5log units or greater change from baseline were nine of 23 (39%) for LNP subjects, and seven of 18 (39%) for LP subjects ( $\chi^2=0.09$ ,  $P=1.0$  [Fisher's exact test]).

On posthoc analysis, a comparison of bifidobacteria and lactobacilli counts between LNP and LP individuals in tests 3 were 10.3log units versus 9.7log units ( $P=0.14$ ), and 11.6log units versus 11.6log units ( $P=0.77$ ), respectively.

In a previous cross-sectional study (15), a very strong agreement (96.7% sensitivity and 92.6% specificity) between genetic classification and the results of the lactose challenge BH<sub>2</sub> tests was found. To explore the suggestion from in vitro data that lactic acid-producing bacteria (bifidobacteria and lactobacilli) may be responsible for reduced BH<sub>2</sub>, the changes

**TABLE 2**  
Counts and bacterial fecal enzymes for both lactase persistent (LP) and lactase nonpersistent (LNP) groups at baseline (test 2) and after 2 weeks of ingesting 25 g of lactose twice a day (test 3)

	LP		LNP	
	Baseline	2-week follow-up	Baseline	2-week follow-up
Bifidobacteria	9.6±0.7	9.7±0.8	9.4±0.7	10.2±1.3
Lactobacilli	11.6±1.0	11.6±0.9	11.0±1.2	11.6±1.1
Total anaerobes	12.2±1.1	12.2±0.9	11.5±1.2	11.9±0.9
Enterobacteriaceae	7.0±1.1	6.7±1.1	6.8±1.2	6.9±1.1
Beta-galactosidase	2.0±0.3	1.9±0.3	1.8±0.4	1.9±0.4
NAG	1.9±0.2	1.8±0.2	1.8±0.2	1.8±0.2

Data presented as mean ± SD. Results are expressed as log<sub>10</sub> colony forming units/g stool for all bacteria, log<sub>10</sub>units/g of stool for beta-galactosidase and log<sub>10</sub> units/mg stool for N-acetyl-beta-D-glucosaminidase (NAG)

in TBH<sub>2</sub> were plotted against the changes in bifidobacteria and lactobacilli in a posthoc analysis using only LNP subjects (Figure 3). The relationship was much stronger for lactobacilli than for bifidobacteria. LP subjects were not included because TBH<sub>2</sub> was essentially zero at baseline and could not decrease further.

To explore whether there were differences in adaptability between LP heterozygotes (CT) and LP homozygotes (TT), the LP group was further investigated with additional posthoc analyses. There were no statistically significant differences in bifidobacteria counts between the CT and TT genotypes.

Table 2 shows the mean bacterial counts and mean fecal enzyme activities for LNP and LP subjects at baseline and follow-up. Changes between these two periods were evaluated in each individual with respect to the secondary comparisons of bacterial counts for Enterobacteriaceae and total anaerobic bacteria, β-gal and NAG stool enzymes, TBH<sub>2</sub> and TSS (LNP versus LP). None of the comparisons achieved statistical significance, although there were the expected reductions in TBH<sub>2</sub> and TSS after lactose challenge (TBH<sub>2</sub>: -57.9 ppm [95% CI -133.4 to 17.7];  $P=0.13$  and TSS: -4.6 units [95% CI -10.0 to 0.8;  $P=0.11$ ]).

The results of the sensitivity analysis for the primary outcome using only complete case scenario data (n=25) found that the differences in the change scores for bifidobacteria was 0.72 (95% CI -0.14 to 1.58;  $P=0.10$ ), and for lactobacilli was 0.08 (95% CI -1.42 to 1.59;  $P=0.91$ ). These results are qualitatively similar to the results of the primary analysis.

## DISCUSSION

The present pilot study supports the notion that prebiotic effects of lactose in humans occur predominantly in LNP rather than in LP individuals, even with high-dose lactose intake. While reduction in TBH<sub>2</sub> following the adaptive period did not achieve the expected results in LNP subjects, a significant change was demonstrated in bifidobacteria in LNP compared with LP participants. The numerically reduced TBH<sub>2</sub> observed in LNP was nevertheless closely linked with lactobacilli.

The concept of a prebiotic, nonabsorbed carbohydrate that selectively alters lower intestinal bacterial flora in a beneficial way to the host was first introduced by Gibson and Roberfroid (32),

and altered to include detection of lactic acid bacteria in stool (33,34). In lactose-intolerant individuals, continued consumption of DFs can lead to adaptation (3,5-8,35-40), but not because of intestinal lactase induction (41) – both bifidobacteria and lactobacilli could be responsible (5,42,43). Also, lactose was shown to induce bifidobacteria proliferation in an in vitro model of the colon (44). Furthermore, lactose forms the backbone of some recognized prebiotics (45,46).

In the only other in vivo human study to evaluate fecal flora, Ito and Kimura (31) showed increased lactobacilli and bifidobacteria after a short exposure to lactose in lactose maldigesters. This study also found reduced bacteroides and *Clostridium* species. In our study, the mean change in bifidobacteria after a median of 15 days of lactose consumption was a 0.83 log<sub>10</sub> units and approximately two-thirds of the subjects in the LNP group achieved 0.5log (three-fold) or greater change. It should be noted that this magnitude of change for bifidobacteria is similar to those reported with oligofructose (a recognized prebiotic) (47,48). Interestingly, reduction in BH<sub>2</sub> was inversely correlated with lactobacilli in the present study. The effect on bifidobacteria in the LNP group was approximately seven-fold greater than that for the LP group, which showed minimal or virtually no change. Taken together, the in vitro studies and human observations provide support for a prebiotic effect of lactose, as redefined more recently (33,34), and a differential bifidogenic effect on LNP compared with LP phenotypes.

At the lactose doses we used (25 g twice a day), the expected spillage of 2 g/day to 4 g/day into the colon of LP individuals (14) may not have affected microflora in two weeks. We also examined whether C/T LP individuals fared any better with microbial effects and found none. These individuals have a 50% reduction in intestinal lactase (17) and might have been expected to spill more lactose than T/T LP individuals.

Several weaknesses in the present study need to be addressed. First, our results should be interpreted in light of the unanticipated reduction in study power to detect a three-fold change in bacterial counts. Nevertheless, the changes in bifidobacteria counts (from follow-up compared with baseline within groups) between LNP and LP subjects showed a greater difference than this. Therefore, the reason why outcome was not more dramatic may be related to the failure to induce complete

adaptation in LNP participants. Because TBH<sub>2</sub> was used as a control evaluating adaptation – a frequently observed phenomenon (5-8,31,35-40) – the question becomes why was this not achieved. Despite an attempt to use standard methodology in ascertaining consumption of lactose (ie, returned empty containers), we failed to properly record returns. Although historically suggested, we remain uncertain whether all participants explicitly followed the instructions. Second, we chose a period of 15 days of lactose consumption because it was previously shown that 16 days was adequate time for full adaptation to occur (5). However, this still may not have been enough time given the above outlined weaknesses. A longer period of lactose consumption, even with reduced adherence might have achieved the expected statistical significance in the reduction of BH<sub>2</sub>. Full adaptation may have enhanced bacterial effects. Finally, we recognize that due to the methods used to assess stool samples, deterioration in colony counts likely occurred (49); however, this was unavoidable, but due to the blinded nature of the study, the order and genetic status of the participants was unknown to those analyzing stool. As such, similar inaccuracies could have been incurred equally by both groups.

## CONCLUSIONS

Despite limitations, the present study lends support to the concept that lactose exerts selective colonic prebiotic effects on LNP versus LP subjects. A bifidogenic effect of lactose is more evident in LNP individuals. Reduced TBH<sub>2</sub> after lactose intervention is closely linked with metabolism by lactobacilli. Future studies should evaluate dose-time effects in these two populations and the specific species of bacteria affected. The contribution of host/intestinal bacterial interactions to disease is becoming more evident in some diseases (50,51). The possible modifying influence on the pathogenesis of lactose/lactase interactions should be taken into consideration.

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